<u>AMENDMENTS</u>

In the Claims

1 A composition comprising a polynucleotide sequence, wherein the 1.(withdrawn) 2 polynucleotide sequence comprises an AIPL1 sequence within the LCA4 region of chromosome 3 17p13 and is selected from the group consisting of a wild-type AIPL1 sequence and a mutant AIPL1 4 sequence. 1 2.(withdrawn) The composition of claim 1, wherein the mutants are selected from the group 2 consisting of Ala336Δ2, Trp278X, Cys239Arg, M79T, L88X, V96I, T124I, P376S, Q163X, A197P, 3 IVS2-2, G262S, R302L, P351D12, Cys42X (TGT -> TGA), Val33ins 8 bp (GTGATCTT), 4 Leu257del 9 bp (CTCCGGCAC) and mixtures and combinations thereof. 1 3.(withdrawn) A protein comprising SEQ. ID. NOs. 72-78 and variants of the protein of 2 SEQ. ID. NO. 72, or a polypeptide expressed by a polynucleotide comprising a nucleotide sequence 3 selected from the group consisting of SEQ. ID NOs. 1-8 or mutants of SEQ. ID. NO. 1 selected from 4 the group consisting of SEQ. ID Nos. 9-41. 1 4.(withdrawn) A purified polynucleotide sequence comprising a sequence selected from the 2 group consisting of SEO ID NOs. 1-71. 1 5.(withdrawn) A retinal disease diagnostic library comprising anti-sense DNA sequences, 2 each sequence corresponding to a DNA sequence including a mutation of the AIPL1 gene selected 3 from the group consisting of SEQ. ID Nos. 9-41 and mixtures and combinations thereof. 1 6.(withdrawn) A primer comprising an AIPL1 sequence, wherein the AIPL1 sequence is 2 selected from the group consisting of a wild-type AIPL1 sequence and a mutant AIPL1 sequence, 3 wherein the mutant-AIPL1 contributes to a retinal disease. 7.(withdrawn)

The primer of claim 6, further comprising a polynucleotide sequence selected

from the group consisting of SEQ ID NOs. 42-47 and 60-71.

1	8.(withdrawn	A probe comprising an AIPL1 sequence, wherein the AIPL1 sequence is			
2	selected from the group consisting of a wild-type AIPL1 sequence and a mutant AIPL1 sequence,				
3	wherein the mutant-AIPL1 contributes to a retinal disease.				
1	9.(original)	A method to determine if an animal has a retinal disease or has a propensity to pass			
2	a retinal disease to offspring, comprising the steps of:				
3	(A)	extracting polynucleotide from a cell or sample;			
4	(B)	determining if the polynucleotide contains a mutation in an AIPL1 encoding or			
5		regulating region; and			
6	(C)	correlating the presence of the mutation as an indication of a retinal disease or a			
7		propensity to pass a retinal disease to offspring.			
1	10.(original)	The method of claim 9, further comprising the steps of:			
2	obtaining a patient sample; and				
3	amplifying the polynucleotide.				
1	11.(original)	The method of claim 10, wherein the amplifying is done via polymerase chain			
2	reaction.				
1	12.(original)	The method of claim 9, wherein the determining is done via polynucleotide sequence.			
1	13.(currently	amended) The method of claim 9, wherein the mutations is are selected from the			
2	group consisting	ng of Ala336Δ2, Trp278X , Cys239Arg, M79T, L88X, V96I, T124I, P376S, Q163X,			
3	A197P, IVS2-2, G262S, R302L, P351D12, Cys42X (TGT -> TGA), Val33ins 8 bp (GTGATCTT),				
4	Leu257del 9 b	p (CTCCGGCAC) and mixtures and combinations thereof.			
1	14.(withdraw	n) A therapeutic method to treat retinal disease comprising the step of			
2	administering to an animal an effective amount of a protein encoded by a wild-type AIPL1 gene or				
3	a polynucleotide sequence a wild-type AIPL1 gene or a retinal medication designed to ameliorate				
4	disease symptoms to the natient if the mutation is detected or mixtures or combinations thereof				

1	15.(withdrawn)	The method of claim 14, wherein the medication is an drug that inhibits retinal					
2	cell death.						
1	16.(withdrawn)	The method of claim 14, wherein the mutations are selected from the group					
2	consisting of Ala33	6Δ2, Trp278X, Cys239Arg, M79T, L88X, V96I, T124I, P376S, Q163X, A197P,					
3	IVS2-2, G262S, R	IVS2-2, G262S, R302L, P351D12, Cys42X (TGT -> TGA), Val33ins 8 bp (GTGATCTT),					
4	Leu257del 9 bp (CTCCGGCAC) and mixtures and combinations thereof.						
1	17.(withdrawn)	A method to determine if a patient has a mutant AIPL1 gene comprising:					
2	(a) extra	acting AIPL1 polypeptide from a cell or sample from the patient;					
3	(B) deter	rmining if the polypeptide contains an AIPL1 mutation; and					
4	(C) corre	elating the mutation as an indication of a retinal disease.					
1	18.(withdrawn)	The method of claim 17, wherein the mutations are selected from the group					
2	consisting of Ala33	6Δ2, Trp278X, Cys239Arg, M79T, L88X, V96I, T124I, P376S, Q163X, A197P,					
3	IVS2-2, G262S, R	302L, P351D12, Cys42X (TGT -> TGA), Val33ins 8 bp (GTGATCTT),					
4	Leu257del 9 bp (C7	Leu257del 9 bp (CTCCGGCAC) and mixtures and combinations thereof.					
1	19.(withdrawn)	A method of producing a cell expressing an AIPL1 mutation comprising					
2	transfecting a cell	with a polynucleotide sequence having at least one AIPL1 mutation in the					
3	sequence.						
1	20.(withdrawn)	The method of claim 19, wherein the encoded mutation is selected from the					
2	group consisting of	group consisting of are selected from the group consisting of Ala336Δ2, Trp278X, Cys239Arg,					
3	M79T, L88X, V96I, T124I, P376S, Q163X, A197P, IVS2-2, G262S, R302L, P351D12, Cys42X						
4	(TGT -> TGA), Val	(TGT -> TGA), Val33ins 8 bp (GTGATCTT), Leu257del 9 bp (CTCCGGCAC) and mixtures and					
5	combinations thereof.						
1	21.(currently ame	nded) A method for determining the presence of an AIPL1 mutant in a					
2	patient sample, whi	ch comprises:					
3	(A) isola	ting polynucleotide extracted from the patient sample;					

4	(B) Hydr	idizing a detectably labeled offgonucleotide to the polynucleotide isolated in step	
5	(b), 1	the oligonucleotide having at its 3' end at least 15 nucleotides complementary to	
6	a wi	ld type polynucleotide sequence having at least one mutation;	
7	(C) atter	npting to extend the oligonucleotide at its 3'-end;	
8	(D) asce	rtaining the presence or absence of a detectably labeled extended	
9	oligo	onucleotide; and	
0	(E) corre	elating the presence or absence of a detectably labeled extended oligonucleotide	
1	in st	ep (e) with the presence or absence of a AIPL1 <u>Trp278X</u> mutation.	
1	22.(original) The	method of claim 21, further comprising taking a patient sample prior to the	
2	isolating step.		
1	23.(original) The	method of claim 21, wherein the isolated nucleic acid is amplified prior to	
2	hybridization.		
1	24.(original) The	method of claim 21, wherein the detectable label on the oligonucleotide is an	
2	enzyme, radioisotop	pe or fluorochrome.	
1	25.(withdrawn)	A test kit useful for the detection of AIPL1 mutations comprising a container	
2	containing at least one polynucleotide capable of hybridizing with a polynucleotide encoding at least		
3	one mutation selected from the group consisting of Ala336Δ2, Trp278X, Cys239Arg, M79T, L88X,		
4	V96I, T124I, P376S, Q163X, A197P, IVS2-2, G262S, R302L, P351D12, Cys42X (TGT -> TGA),		
5	Val33ins 8 bp (GTGATCTT), Leu257del 9 bp (CTCCGGCAC) and mixtures and combinations		
6	thereof.		
1	26.(withdrawn)	A method of screening compounds to determine their effectiveness in	
2	counteracting a cell	's retinal behavior due to a mutation in its AIPL1 gene comprising:	
3	(A) conta	acting the compound with a cell including a mutation is its AIPL1 gene where	
4	the	mutation is selected from the group consisting of Ala336Δ2, Trp278X,	
5	Cvs?	239Arg, M79T, L88X, V96I, T124I, P376S, O163X, A197P, IVS2-2, G262S	

0		K302L, P351	D12, Cys42X (1G1 -> 1GA), Val33ins 8 bp (G1GA1C11), Leu25 /del	
7		9 bp (CTCC	GGCAC) and mixtures and combinations thereof; and	
8	(B)	determining if the cell is affected by the compound.		
1	27.(currently	amended)	A method to determine if a cell or sample has an AIPL1 mutation	
2	comprising:			
3	(A)	extracting po	olynucleotide from a cell;	
4	(B)	amplifying polynucleotides which encode AIPL1; and		
5	(C)	determining if the polynucleotide contains a <u>Trp278X</u> mutation;		
6	(D)	correlating the presence of the mutation as an indication of a retinal disease or a		
7		propensity to	pass a retinal disease to offspring.	